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To: Chancellor, Erin [chancellor.erin@epa.gov]

Subject: Ethylene oxide key findings

Hey Erin,

I will go ahead and share our key findings with you, but please hold it close for now. The wording on this is subject to change in our public comment draft, but the conclusions should not change. The important things are highlighted below in yellow.

Basically, we plugged the data used to derive the EPA model back into the model to see how many cancers the model predicts are in the data used to derive the model. Out of 17+k occupationally exposed workers, 53 developed lymphoid cancer. When you plug the exposure data back into EPA's model, it predicts that 1,179 workers developed cancer, way higher than the actual 53. Our model predicts that 59 developed cancer, just slightly higher than the actual 53. So our model is more predictive. EPA's model is over predictive. Though I know NCEA and IRIS will not want to budge off of their position, it's kind of hard to argue about the math. It's pretty straightforward.

Like I said, we're cleaning up the document and making sure we are precise in our wording, but should have it out for public comment by the end of next week at the latest.

Best, Mike

Key Findings

- Ethylene oxide (EtO) is a chemical with many industrial applications, with particular use as a sterilant for medical devices.
- Because EtO is emitted in Texas and has been determined to be a carcinogen, the TCEQ undertook a carcinogenic dose-response assessment and derivation of an effect screening level for this chemical.
- Review of the EtO literature demonstrated that EtO operates by a direct-acting mutagenic mode of action (MOA) and suggests that the EtO cancer dose-response should be no more than linear overall with sublinearity expected by both TCEQ and USEPA at endogenous levels and below.
- In addition, EtO is produced endogenously, and an ambient air concentration of ~1.3 ppb would be required to increase the internal dose (endogenous + exogenous) of EtO by 1 standard deviation. Therefore, ambient EtO concentrations of less than 1 ppb would not be expected to produce substantively more risk than endogenously-produced EtO.
- Consistent with TCEQ guidelines (TCEQ 2015), we reviewed recently derived toxicity factors and guideline air levels to determine if there is a toxicity factor or guideline air level that is suitable for adoption by the TCEQ. As such, we reviewed the USEPA's recently completed Evaluation of the Inhalation Carcinogenicity of Ethylene Oxide (USEPA 2016). The USEPA derived a unit risk factor (URF) of 7.1E-3 per ppb, which corresponds to a 1 in 100,000 excess cancer risk air concentration of 0.001 ppb.
- The human data available for deriving the USEPA's EtO toxicity factor came from two very high exposure occupational cohorts (UCC and NIOSH) that provide no information about the shape of the dose-response curve at low (i.e., environmentally-relevant) EtO concentrations. The TCEQ agrees with

USEPA's determination that in the low-dose range a sublinear dose-response is "highly plausible," based on the MOA and information about endogenous production of EtO.

- In contrast to their determination that the low-dose region of the EtO dose-response curve is highly plausibly sublinear, USEPA ultimately chose to model EtO-induced lymphoid cancer with an overall supra-linear two-piece spline model that has a very steep linear slope in the low-dose region.
- The TCEQ evaluated USEPA's URF and overall supra-linear (i.e., linear two-piece spline) modeling choice in the context of the available observed data to determine the validity of the modeling and URF:

 - O Population-Level Lymphoid Cancer Risk Using measured concentrations of a biomarker of internal EtO exposure (an EtO-specific protein adduct in blood), it can be estimated that the mean amounts of endogenous EtO levels would be equivalent to ambient concentrations of EtO of 1.9 ppb in non-smokers and 18.8 ppb in smokers. Accordingly, at measured endogenous levels of EtO, the USEPA's URF would predict a population-wide lymphoid cancer rate of 3.8% (in the absence of any exogenous EtO exposure or other potential causes of lymphoid cancer). By contrast, the USEPA-cited lymphoid cancer background rate (which would have many contributing factors, not just a single chemical) is 3%, demonstrating that USEPA's URF overestimates observable lymphoid cancer risk based on endogenous levels alone.
 - Lymphoid Cancer Risk from Cohort Studies The UCC cohort shows no statistically significant increased risk of lymphoid cancer with EtO exposure. The NIOSH cohort shows statistically significant increased risk of lymphoid cancer only with the highest EtO cumulative exposure quintile. These data are not consistent with USEPA's selected overall supra-linear two-piece spline model because that model would predict statistically increased risks at lower EtO cumulative exposures, not just at the highest (see below).
 - Model Fit with Observed Data USEPA conducted their EtO cancer dose-response modeling using the NIOSH cohort data. To verify that USEPA's final selected model assessment (i.e., upper bound on the linear two-piece spline model) properly fit the original data, we used it to predict the expected number of lymphoid cancers based on the same NIOSH individual exposure data as USEPA used for modeling. Whereas 53 lymphoid cancer deaths were observed in this cohort of 17,530 workers, USEPA's selected dose-response model assessment predicted 1,179 (901, 1574) lymphoid cancer deaths in this same cohort. Similarly, USEPA's final selected model assessment statistically significantly over-predicts lymphoid cancer deaths in every cumulative exposure quintile and indicates that statistically increased lymphoid cancer should have occurred in every exposure quintile (including the lowest), when in fact this did not occur. This demonstrates unequivocally that USEPA's selected model assessment cannot be validated by the data that was used to derive it, and this model is not appropriate to use for estimates of population risk.
- The TCEQ determined that USEPA's use of an overall supra-linear dose-response model (i.e., the upper bound of the linear two-piece spline model) to derive their URF: 1) is not justified by the MOA data (which support a no-more-than linear dose-response); 2) is not consistent with predicted population risk from endogenous EtO for lymphoid cancer; and 3) grossly over-estimates the number of lymphoid cancers in the cohort from which the dose-response model was derived. Therefore, the TCEQ found that USEPA's EtO inhalation URF lacks scientific credibility and the TCEQ did not adopt it for this evaluation.

- The TCEQ conducted a systematic review for studies that could inform the derivation of a cancer URF for inhalation exposures to EtO. This review identified key epidemiological data from two cohorts of occupationally-exposed workers, and Cox proportional hazards modeling was conducted to model the EtO-cancer dose-response.
- The TCEQ ultimately chose lymphoid cancer as the critical cancer endpoint, using a 15-year EtO exposure lag with results for NIOSH males being more conservative, to calculate a **URF of 2.5E-6 per ppb (1.4E-6 per ug/m³)** and a ^{chronic}ESL_{nonthreshold(c)} of **4 ppb (7 ug/m³)** at an excess cancer risk level of 1 in 100,000.
- As with USEPA's URF, we evaluated the TCEQ's URF in the context of the available observed data to determine the validity of the modeling and URF:
 - o <u>Endogenous levels of EtO</u> Compared to endogenous EtO levels, the TCEQ's ESL of 4 ppb would produce an internal exposure equivalent to between the 90th-95th percentile of the normal endogenous range and could biologically plausibly be associated with excess risk above and distinguishable from the endogenous levels.
 - Population-Level Lymphoid Cancer Risk At measured endogenous levels of EtO, the TCEQ's URF would predict a population-wide lymphoid cancer rate that is lower than the cited background population cancer rate of 3%.
 - <u>Lymphoid Cancer Risk from Cohort Studies</u> The standard Cox proportional hazards model of lymphoid cancer did not show a relationship with EtO exposure that was statistically significantly different from zero. Therefore, by assuming a significant positive slope in the EtOcancer association, the TCEQ is making a conservative decision to assume that EtO is causing lymphoid cancer in the exposed workers in the NIOSH cohort.
 - Model Fit with Observed Data To verify that the TCEQ's model properly fit the original data, we calculated the expected number of lymphoid cancers based on the individual exposure estimates for the NIOSH cohort (also used by USEPA). Whereas 53 lymphoid cancer deaths were observed in this cohort of 17,530 workers, the TCEQ's selected dose-response assessment (i.e., upper bound of the Cox proportional hazard model) predicted 59 (45, 78) lymphoid cancer deaths. Similarly, TCEQ's selected assessment neither significantly over- or under-estimated lymphoid cancer for any exposure quintile. This demonstrates that the TCEQ's model selection provides a good fit to the observed number of lymphoid cancer deaths in the NIOSH cohort.
- The TCEQ determined that the use of Cox proportional hazards models to derive a URF for inhalation EtO cancer risk: 1) is justified by the MOA data showing EtO to be a direct-acting carcinogen whose effects, particularly at doses near the endogenous range, would be buffered by cellular repair mechanisms; 2) is consistent with population background risk level considering background endogenous EtO levels (i.e., does not overestimate population risks for lymphoid cancer); and 3) accurately estimates the number of lymphoid cancers in the cohort from which the dose-response model was derived. Therefore, the TCEQ's EtO URF has a sound scientific basis and will be adopted for review of air concentration data and for use in air permit reviews.



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